

Clarius Mobile Health Corp. % Ms. Agatha Szeliga Director, Regulatory Affairs 130-2985 Virtual Way Vancouver, British Columbia V5M 4X7 CANADA

Re: K213436 November 15, 2021

Trade/Device Name: Clarius Ultrasound Scanner

Regulation Number: 21 CFR 892.1550

Regulation Name: Ultrasonic pulsed doppler imaging system

Regulatory Class: Class II

Product Code: IYN Dated: October 21, 2021 Received: October 22, 2021

Dear Ms. Szeliga:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. Although this letter refers to your product as a device, please be aware that some cleared products may instead be combination products. The 510(k) Premarket Notification Database located at https://www.accessdata.fda.gov/scripts/cdrh/cfdocs/cfpmn/pmn.cfm identifies combination product submissions. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801 and Part 809); medical device reporting (reporting of medical device-related adverse events) (21 CFR

803) for devices or postmarketing safety reporting (21 CFR 4, Subpart B) for combination products (see https://www.fda.gov/combination-products/guidance-regulatory-information/postmarketing-safety-reporting-combination-products); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820) for devices or current good manufacturing practices (21 CFR 4, Subpart A) for combination products; and, if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to https://www.fda.gov/medical-device-problems.

For comprehensive regulatory information about medical devices and radiation-emitting products, including information about labeling regulations, please see Device Advice (https://www.fda.gov/training-and-continuing-education/cdrh-learn) and CDRH Learn (https://www.fda.gov/training-and-continuing-education/cdrh-learn). Additionally, you may contact the Division of Industry and Consumer Education (DICE) to ask a question about a specific regulatory topic. See the DICE website (https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice">https://www.fda.gov/medical-devices/device-advice-comprehensive-regulatory-assistance/contact-us-division-industry-and-consumer-education-dice) for more information or contact DICE by email (DICE@fda.hhs.gov) or phone (1-800-638-2041 or 301-796-7100).

Sincerely,

For

Thalia T. Mills, Ph.D.

Director
Division of Radiological Health
OHT7: Office of In Vitro Diagnostics
and Radiological Health
Office of Product Evaluation and Quality
Center for Devices and Radiological Health

Enclosure

DEPARTMENT OF HEALTH AND HUMAN SERVICES Food and Drug Administration

Indications for Use

Type of Use (Select one or both, as applicable)

Prescription Use (Part 21 CFR 801 Subpart D)

Form Approved: OMB No. 0910-0120
Expiration Date: 06/30/2023

Expiration Date: 06/30/2023
See PRA Statement below.

510(k) Number <i>(if known)</i>
K213436
Device Name
Clarius Ultrasound Scanner
Indications for Use (Describe)
The Clarius Ultrasound Scanner is a software-based ultrasound imaging system and accessories, intended for diagnostic
imaging. It is indicated for diagnostic ultrasound imaging and fluid flow analysis in the following applications: ophthalmic, fetal, abdominal, intra-operative (non-neurological), pediatric, small organ, cephalic (adult), trans-rectal, trans-vaginal, musculo-skeletal (conventional, superficial), urology, gynecology, cardiac (adult, pediatric), peripheral vessel, carotid, and procedural guidance of needles into the body.
The system is a transportable ultrasound system intended for use in environments where healthcare is provided by trained healthcare professionals.

CONTINUE ON A SEPARATE PAGE IF NEEDED.

Over-The-Counter Use (21 CFR 801 Subpart C)

This section applies only to requirements of the Paperwork Reduction Act of 1995.

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Department of Health and Human Services Food and Drug Administration Office of Chief Information Officer Paperwork Reduction Act (PRA) Staff PRAStaff@fda.hhs.gov

"An agency may not conduct or sponsor, and a person is not required to respond to, a collection of information unless it displays a currently valid OMB number."

510(k) Summary - K213436

This 510(k) summary of safety and effectiveness information is submitted in accordance with the requirements of 21 CFR § 807.92.

Subject Device Trade Name: Clarius Ultrasound Scanner

Subject Device Model Numbers: C3 HD3, C7 HD3, EC7 HD3, L7 HD3, L15 HD3, PA HD3, L20 HD3

<u>Common Name:</u> Diagnostic Ultrasound System and Accessories

Regulation Number, Name and Product Codes:

Regulation Number	Regulation Name	Product Code
21 CFR § 892.1550	Ultrasonic Pulsed Doppler Imaging System	IYN
21 CFR § 892.1560	Ultrasound Pulsed Echo Imaging System	IYO
21 CFR § 892.1570	Diagnostic Ultrasonic Transducer	ITX

FDA 510(k) Review Panel: Radiology

<u>Classification:</u> Class II

Manufacturer: Clarius Mobile Health Corp.

130-2985 Virtual Way

Vancouver, BC V5M 4X7 Canada

Contact Name: Agatha Szeliga

Director, Regulatory Affairs agatha.szeliga@clarius.com

<u>Date 510(k) Summary Prepared:</u> October 20, 2021

Predicate Device Information:

Device Trade Name:	Clarius Ultrasound Scanner
510(k) Reference:	K192107
Submitter Name:	Clarius Mobile Health Corp.
Regulation Name:	Ultrasonic Pulsed Doppler Imaging System
Classification Product Code(s):	IYN
Subsequent Product Codes	IYO, ITX
Regulation Number:	21 CFR § 892.1550; 21 CFR § 892.1560; 21 CFR §
	892.1570
Classification:	Class II

Note: The predicate device has not been subject to a design-related recall.

Device Description

The Clarius Ultrasound Scanner is a portable, general-purpose, software-controlled, diagnostic ultrasound system used to acquire and display high-resolution, real-time ultrasound data through an off-the-shelf (OTS) iOS or Android device. The Clarius Ultrasound Scanner comprises a series of wireless transducers employing Bluetooth and Wi-Fi-based technology to communicate with traditional tablet/smartphone devices via direct Wi-Fi. This allows the user to export ultrasound images and display them across a range of portable personal devices.

The system is a transportable ultrasound system intended for use in environments where healthcare is provided by trained healthcare professionals, including the emergency medical services (EMS) environment.

The Clarius Ultrasound Scanner is intended for use by qualified healthcare practitioners (e.g., doctors, nurses, sonographers) who are trained in the use of ultrasound imaging technology.

The Clarius Ultrasound Scanner comprises the following:

Transducers/ Scanners (various models)	C3 HD3, C7 HD3, EC7 HD3, L7 HD3, L15 HD3, PA HD3,
Transducers/ Scanners (various moders)	L20 HD3
Coffee	Clarius Ultrasound App (Clarius App) for iOS;
Software	Clarius Ultrasound App (Clarius App) for Android
Accession	Clarius Charger
Accessories	Clarius Fan

Indications for Use for the Clarius Ultrasound Scanner

The Clarius Ultrasound Scanner is a software-based ultrasound imaging system and accessories, intended for diagnostic imaging. It is indicated for diagnostic ultrasound imaging and fluid flow analysis in the following applications: ophthalmic, fetal, abdominal, intra-operative (non-neurological), pediatric, small organ, cephalic (adult), trans-rectal, trans-vaginal, musculo-skeletal (conventional, superficial), urology, gynecology, cardiac (adult, pediatric), peripheral vessel, carotid, and procedural guidance of needles into the body.

The system is a transportable ultrasound system intended for use in environments where healthcare is provided by trained healthcare professionals.

Comparison of the Subject Device and Predicate Device for Demonstration of Substantial Equivalence

Criteria	SUBJECT DEVICE	PREDICATE DEVICE
	Clarius Ultrasound Scanner	Clarius Ultrasound Scanner
510(k) Holder/	Clarius Mobile Health Corp.	Clarius Mobile Health Corp.
Manufacturer		
Submission	Current Submission	K192107
Reference		
510(k) Track	Track 3	Track 3
Product Codes	IYN ¹ , IYO ² , ITX ³	IYN ¹ , IYO ² , ITX ³

Criteria	SUBJECT DEVICE	PREDICATE DEVICE
	Clarius Ultrasound Scanner	Clarius Ultrasound Scanner
Regulation	21 CFR 892.1550 ¹	21 CFR 892.1550 ¹
Number	21 CFR 892.1560 ²	21 CFR 892.1560 ²
	21 CFR 892.1570 ³	21 CFR 892.1570 ³
Regulation Name	Ultrasonic Pulsed Doppler Imaging System ¹ ;	Ultrasonic Pulsed Doppler Imaging System ¹ ;
	Ultrasonic Pulsed Echo Imaging System ² ;	Ultrasonic Pulsed Echo Imaging System ² ;
	Diagnostic Ultrasonic Transducer ³	Diagnostic Ultrasonic Transducer ³
Transducer	C3 HD3, C7 HD3, EC7 HD3, L7 HD3, L15 HD3,	C3 HD, C7 HD, EC7 HD, L7 HD, L15 HD, PA HD,
Models	PA HD3, L20 HD3	L20 HD
Transducer Types	Convex Array	Convex Array
	Linear Array Phased Array	Linear Array Phased Array
	Intracavity	Intracavity
Intended Use	Diagnostic ultrasound imaging and fluid flow	Diagnostic ultrasound imaging and fluid flow
intended 03e	analysis of the human body	analysis of the human body
Indications for	The Clarius Ultrasound Scanner is a software-	The Clarius Ultrasound Scanner is a software-
Use and Clinical	based ultrasound imaging system and	based ultrasound imaging system and
Usage	accessories, intended for diagnostic imaging.	accessories, intended for diagnostic imaging.
	It is indicated for diagnostic ultrasound	It is indicated for diagnostic ultrasound
	imaging and fluid flow analysis in the	imaging and fluid flow analysis in the
	following applications: ophthalmic, fetal,	following applications: ophthalmic, fetal,
	abdominal, intra-operative (non-	abdominal, intra-operative (non-
	neurological), pediatric, small organ, cephalic	neurological), pediatric, small organ, cephalic
	(adult), trans-rectal, trans-vaginal, musculo-	(adult), trans-rectal, trans-vaginal, musculo-
	skeletal (conventional, superficial), urology,	skeletal (conventional, superficial), urology,
	gynecology, cardiac (adult, pediatric),	gynecology, cardiac (adult, pediatric),
	peripheral vessel, carotid, and procedural	peripheral vessel, carotid, and procedural
	guidance of needles into the body.	guidance of needles into the body.
	The system is a transportable ultrasound	The system is a transportable ultrasound
	system intended for use in environments	system intended for use in environments
	where healthcare is provided by trained	where healthcare is provided by trained
	healthcare professionals.	healthcare professionals.
	Ophthalmic	Ophthalmic
	Fetal	• Fetal
	Abdominal	Abdominal
	Intraoperative (Ab/Vasc)	Intraoperative (Ab/Vasc)
	Pediatric	Pediatric
	Small Organ	Small Organ
	Adult cephalic	Adult cephalic
	Trans-rectal	Trans-rectal
	Trans-vaginal	Trans-vaginal
	Musculoskeletal (conventional)	Musculoskeletal (conventional)
	Musculoskeletal (superficial)	Musculoskeletal (superficial)
	Urology	Urology
	Gynecology	Gynecology
	Cardiac adult	Cardiac adult

Criteria	SUBJECT DEVICE	PREDICATE DEVICE
	Clarius Ultrasound Scanner	Clarius Ultrasound Scanner
	Cardiac pediatric	Cardiac pediatric
	Peripheral vessel	Peripheral vessel
	Carotid	Carotid
	Needle guidance	Needle guidance
Principle of	Piezoelectric material in the system's	Piezoelectric material in the system's
Operation	transducer transmits high frequency, non-	transducer transmits high frequency, non-
	ionizing sound waves to the designated	ionizing sound waves to the designated
	region of the body and converts the	region of the body and converts the
	subsequent echoes detected to electronic	subsequent echoes detected to electronic
	signals in order to construct an image of the	signals in order to construct an image of the
	internal structures of an anatomical field. The	internal structures of an anatomical field. The
	image is sent wirelessly from the transducer	image is sent wirelessly from the transducer
	to an external iOS or Android viewing device	to an external iOS or Android viewing device
	on which the image can be displayed.	on which the image can be displayed.
Power Source	Internal integrated built-in (non-removable)	Removable lithium-ion battery
	lithium-ion battery	·
Display	iOS or Android mobile device	iOS or Android mobile device
Wireless	Communicates wirelessly via Wi-Fi and	Communicates wirelessly via Wi-Fi and
Capability	Bluetooth	Bluetooth
Portability	Portable ultrasound system	Portable ultrasound system
System	Transducers/scanners	Transducers/scanners
Components	Software (Clarius App)	Software (Clarius App)
	Accessories (Charger and Fan)	Accessories (Charger and Fan)
Modes of	B-mode	B-mode
Operation	M-mode	M-mode
	Color Doppler	Color Doppler
	Power Doppler	Power Doppler
	Pulse-Wave Doppler (PWD)	Pulse-Wave Doppler (PWD)
	Combined (B+M; B+CD; B+PD; B+PWD)	Combined (B+M; B+CD; B+PD; B+PWD)
Safety Standards	The Clarius Ultrasound Scanner complies with	The Clarius Ultrasound Scanner complies with
,	the following safety standards:	the following safety standards:
	60601-1	60601-1
	60601-1-2	60601-1-2
	60601-1-6	60601-1-6
	60601-1-12	60601-1-12
	60601-2-37	60601-2-37

Non-Clinical Performance Testing of the Clarius Ultrasound Scanner

The HD3 scanners of the Clarius Ultrasound Scanner family of devices were designed and developed by Clarius Mobile Health Corp. in accordance with the applicable requirements and standards to establish performance and safety of the device. The device's safety and performance were verified by tests conducted by Clarius and accredited third-party laboratories. Validation testing was performed to ensure that the final product is capable of meeting the requirements for the specified clinical applications and performs as intended to meet users' needs, while demonstrating substantial equivalence to the predicate device.

Non-clinical performance testing of the HD3 transducers of the Clarius Ultrasound Scanner device family demonstrate compliance to the following standards:

Standard	Title of Standard
IEC 60601-1:2012	Medical electrical equipment – Part 1: General requirements for basic safety
	and essential performance
IEC 60601-1-2:2014	Medical electrical equipment – Part 1-2: General requirements for basic safety
	and essential performance – Collateral Standard: Electromagnetic
	Compatibility – Requirements and tests
IEC 60601-1-6 Edition 3.1 2013-	Medical electrical equipment - Part 1-6: General Requirements for Basic Safety
10	and Essential Performance - Collateral Standard: Usability
IEC 60601-1-12 Edition 1.0 2014-	Medical electrical equipment - Part 1-12: General Requirements for Basic
06	Safety and Essential Performance - Collateral Standard: Requirements for
	Medical Electrical Equipment and Medical Electrical Systems Intended for Use
	in the Emergency Medical Services Environment
IEC 60601-2-37:2015	Medical electrical equipment – Part 2-37: Particular requirements for the basic
	safety and essential performance of ultrasonic medical diagnostic and
	monitoring equipment
ISO 10993-1:2018	Biological evaluation of medical devices – Part 1: Evaluation and testing within
	a risk management process
AIUM/NEMA UD 2-2004	NEMA Standards Publication UD 2-2004 (R2009) Acoustic Output
	Measurement Standard for Diagnostic Ultrasound Equipment Revision 3.
	(Radiology)
AIUM/NEMA UD 3-2004	NEMA Standards Publication UD 3-2004 (R2009) Standard for Real-Time
	Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic
	Ultrasound Equipment
IEC 62304:2006/AMD 1:2015	Medical device software – Software life cycle processes
IEC 62366-1:2015	Medical devices – Application of usability engineering to medical devices
ISO 15223-1:2016	Medical devices — Symbols to be used with medical device labels, labelling
	and information to be supplied
ISO 14971:2019	Medical Devices – Application of Risk Management to Medical Devices
IEC 60529:2013	Degrees of protection provided by enclosures (IP Code)
IEC 62133:2012	Secondary cells and batteries containing alkaline or other non-acid electrolytes
	- Safety requirements for portable sealed secondary cells, and for batteries
	made from them, for use in portable applications
IEC 61157:2013	IEC 61157: Standard means for the reporting of the acoustic output of medical
	diagnostic ultrasonic equipment

Conclusion & Summary of Substantial Equivalence

Based on the information presented in this Special 510(k) premarket notification, and based on the fundamental scientific technology, technological characteristics, principle of operation, intended use, environment of use, and indications for use, the modified Clarius Ultrasound Scanner (HD3 scanners) has been determined to be substantially equivalent in terms of safety and effectiveness to the predicate device, the Clarius Ultrasound Scanner (510(k)-cleared in K192107).

The differences in design between the subject device and the predicate device do not raise any issues related to safety or effectiveness.